

Amendments to the Claims:

This listing of claims will replace all prior versions, and listings, of claims in the application:

Listing of Claims:

Claim 1. (original): Solid pharmaceutical composition comprising

- (a) an effective amount of ramipril and/or a pharmaceutical acceptable salt thereof and
- (b) one or more pharmaceutically acceptable excipients,

characterized in that the composition is stabilized by having a suitably low water content of less than about 4.0 weight-% measured by loss-on-drying or of less than about 5.5 weight-% measured by Karl-Fischer-analysis.

Claim 2. (original): Composition according to claim 1, wherein the water content is less than about 4.5 weight-% measured by Karl-Fischer-analysis.

Claim 3. (original): Composition according to claim 1, wherein the water content is less than about 3.0 weight-% measured by loss-on-drying.

Claim 4. (currently amended): Composition according to ~~any of the preceding claims~~ 1, wherein ramipril and/or a pharmaceutical acceptable salt thereof is in form of pharmaceutically acceptable anhydrate, solvate and/or, hydrate and/or in crystalline and amorphous form.

Claim 5. (currently amended): Composition according ~~any of the preceding claims~~ 1, wherein the pharmaceutical composition is a tablet.

Claim 6. (currently amended): Composition according to claim 5, wherein the tablet is ~~suitably~~ coated to generate a ~~filmcoated~~ film coated tablet and/or a pill.

Claim 7. (currently amended): Composition according to claim 1[[- 4]], wherein the pharmaceutical composition is a capsule.

Claim 8. (currently amended): Composition according to claim 1[[- 4]], wherein the pharmaceutical composition is a sachet.

Claim 9. (currently amended): Composition according to ~~any of the preceding claims~~ 1, wherein the excipients have a suitably low water content.

Claim 10. (original): Composition according to claim 9, wherein one of said excipients is microcrystalline cellulose.

Claim 11. (currently amended): Composition according to claim 1[[- 9]], wherein ~~one of said excipients is~~ are selected from the group consisting of Avicel PH 112, starch, Starch 1500 LM,

silicon dioxide, Sylloid AL-1 FP, calcium hydrogen phosphate, Dicafos A or A Tab, anhydrous Emcompress, lactose, Pharmatose DCL 21, mannitol, Perlitol, calcium sulphate, Destab, Drierte, and mixtures thereof.

Claims 12-23 (canceled).

Claim 24. (currently amended): Composition according to ~~any of the preceding claims 1~~ where one or more excipients are dried prior to use or throughout the manufacturing process to achieve the required level of water content.

Claim 25. (currently amended): Process for the preparation of a composition according to ~~any of the preceding claims 1~~, wherein environmental conditions during manufacture are maintained at a relative humidity equal or less than 35% at ambient temperature.

Claim 26. (currently amended): Process for the preparation of a composition according to claim 1[[- 23]], wherein environmental conditions during manufacture are maintained at a relative humidity equal or less than 35% ~~at equal or less than 30° C.~~

Claim 27. (currently amended): Process according to ~~any of the preceding claims 1~~, wherein the pharmaceutical composition is packaged into a packaging material suitably tight against penetration of humidity.

Claim 28. (original): Process according to claim 27, wherein the packaging material is a container including lid composed of polyethylene and/or polypropylene and/or glass.

Claim 29. (original): Process according to claim 27, wherein the packaging material is a strip or blister pack composed of aluminium which might be suitably coated or high density polyethylene.

Claim 30. (currently amended): Package comprising a composition according to claims 1[[- 23]] packaged with packaging material suitably tight against penetration of humidity.

Claim 31. (original): Package according to claim 30, wherein the packaging material is a container including lid composed of polyethylene and/or polypropylene and/or glass.

Claim 32. (original): Package according to claim 30, wherein the packaging material is a strip or blister pack composed of aluminium which might be suitably coated or high density polyethylene.